Current Clinical Program Portfolio Oct-17

Award							Funding	Therapeutic				Percent
Number, PI,					General Disease	General Class	(ICOC	Cell (for Cell				Time Into
Institution	Program	Trial Stage	Indication	Therapeutic	Area	of Approach	Approved)	Therapy)	Cell Source	Rationale	Project Goal	Award
NEURO THERAPE	UTICS											
Neurologic Disor	ders: Injuries											
SP3A-07552										Up to 12,000 Americans suffer a spinal cord injury		
Lebkowski,										each year. Leads to a high level of permanent		
Asterias	Strategic			Allogeneic oligodendrocyte				Oligodendrocyte		disability and decreased life expectancy. Currently	Safety. Dosing. Efficacy -	
Biotherapeutics	Partnership III	Ph 1/2a	Spinal Cord Injury	progenitors	Neurologic Injury	Cell Therapy	\$14,323,318	Progenitors	Allogeneic	no approved therapies.	motor improvement.	
										Stroke is a major cause of long-term disability and		
										there are no proven medical treatments for chronic stroke. Intracerebral delivery of modified MSCs	compared to sham surgery	
										provides a well tolerated treament with the	- improvement in motor	
CLIN2-10344	Clinical Trial			Modified bone marrow-derived						potential to improve motor function in these	activity on stroke affected	
Bates, SanBio	Stage Projects	Ph 2b	Ischemic Stroke	mesenchymal stem cells (MSCs)	Neurologic Injury	Cell Therapy	\$19,998,580	MSC	Allogeneic	patients	side.	
, , , , , , , , , , , , , , , , , , , ,				,	, ,	, ,	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			Stroke is the leading cause of adult disability. There		
										is no medical therapy that promotes stroke		
CLIN1-09433	Late Stage									recovery. Cells derived from H9 ESC act via		
Steinberg,	Preclinical			H9 ESC-derived neural stem				NSC or NPC (ESC-		secretion of paracrine factors to modulate brain		
Stanford	Projects	IND	Ischemic Stroke	cells	Neurologic Injury	Cell Therapy	\$5,300,000	derived)	Allogeneic	repair processes in preclinical stroke models.	Obtain an active IND	
Neurologic Disor	ders: Neurodege	nerative				1					1	
	Disease Team									ALS is a devastating disease with no cure. This cell		
DR2A-05320,										therapy intends to support sick motor neurons via		
CLIN2-09284	Therapy Development,			Allogeneic neural progenitor		Genetically				astrocyte replacement and pro-survival growth factors. Allogeneic neural stem cells, genetically		
Svendsen,	Clinical Trial		ALS (Amyotrophic	- ' -	Neurodegenerative	Modified Cell	\$17,842,617,			modified to express GDNF, injected into the spinal	Safety. Dosing. Efficacy -	
Cedars-Sinai	Stage Projects	Ph 1/2a	lateral sclerosis)	GDNF	Disorder	Therapy	\$6,154,067	NSC or NPC	Allogeneic	cord.	Lower limb strength	
		,	, , , , , , , , , , , , , , , , , , , ,			,	<i>+</i> 1,22 1,221			ALS is a fatal neurodegenerative disease for which		T
										there is currently no adequate treatment.		
										Autologous MSCs are propagated ex vivo and		
										induced to secrete neurotrophic factors. NurOwn		
				Autologous MSCs cultured to						cells are returned to the patients in the target area		
CLIN2-09894	Clinical Trial			enhance secretion of growth	Neurodegenerative					of damage. Previous trials showed safety and	Safety and efficacy of	
Kern, Brainstorm	Stage Projects	Ph 3	lateral sclerosis)	factors (NurOwn)	Disorder	Cell Therapy	\$15,912,390	MSC	Autologous	encouraging signs of efficacy.	three repeated doses.	
Eye Disease	I		T .			T .				A	1	
										Age-related macular degeneration is a progressive disease resulting in death of the retinal pigment		
										epithelium (RPE) causing distortion to central vision		
	Duane Roth									and eventually to legal blindness. Incidence -		
	Disease Team			Allogeneic functionally						1:1359 in the US. Approach is replacement therapy	Safety Efficacy - slow	
	Therapy			polarized hESC-derived RPE						with viable RPE cells delivered on a synthetic	disease progression,	
DR3-07438	Development		Adult Macular	monolayers on synthetic		Cell Therapy,				membrane mimicking native state with RPE cells on		
Humayun, USC	III	Ph 1	Degeneration	substrate	Eye Disease	Combination	\$18,922,665	RPE	Allogeneic	Bruch's membrane.	acuity	
										Retinitis pigmentosa (RP) is a progressive retinal		
										degeneration that affects over 1.5 million people		
										worldwide. Unfortunately, treatment is still rather		
										limited. A single sub-retinal injection of human		
										neural progenitor cells offers dramatic preservation		
1604 0025										of vision. Grafted Cells survive for an extended		
LSP1-0835	Late Stage		D-+ii+i-	C. barriard interstance for						period, secrete pro-survival factors and		
Wang, Cedars- Sinai	Preclinical Projects	IND	Retinitis Pigmentosa	Subretinal injection of human neural progenitor cells	Eye Disease	Cell Therapy	\$4,954,514	NPC	Allogeneic	extracellular matrix, reduce oxidative stress response and preserve vision and RPE integrity.	Obtain an active IND	
Jillai	FTOJECIS	טוווט	Figurenitosa	neurai progenitor cens	Lye Disease	сен ппетару	14ر+ردر+ب	INFC	Allogerielt	Retinitis pigmentosa (RP) is a severe form of	Obtain all active IND	
										blindness that runs in families with an incidence of		
										1:4000. Good target for stem cell therapy due to		
DR2A-05739	Disease Team									the defined loss of specific cells. Proposed		
Klassen, UC	Therapy	IND, Ph	Retinitis	Allogeneic retinal progenitor						mechanism: Rescue the light sensing	Safety and efficacy - visual	
Irvine	Development	1/2a	Pigmentosa	cells	Eye Disease	Cell Therapy	\$17,306,668	RPC	Allogeneic	photoreceptors.	acuity.	
											Safety and efficacy -	
CLIN2-09698	Clinical Trial		Retinitis	Allogeneic retinal progenitor						Follow-on study based on Phase 1/2a clinical trial.	improvement in visual	
Klassen, Jcyte	Stage Projects	Ph 2b	Pigmentosa	cells	Eye Disease	Cell Therapy	\$8,295,750	RPC	Allogeneic	Continue to assess safety and establish efficacy.	function at 12 months.	

Award							Funding	Therapeutic			October 26th,	Percent
Number, PI,					General Disease	General Class	(ICOC	Cell (for Cell				Time Into
Institution	Drogram	Trial Stage	Indication	Thoronoutic				-	Call Saurea	Pationalo	Project Goal	
institution	Program	Trial Stage	indication	Therapeutic	Area	of Approach	Approved)	Therapy)	Cell Source		Project Goal	Award
										Limbal stem cell deficiency results in inability to		
										heal following ocular surface injury leading to		
										corneal opacity. Cultivated autologous limbal stem		
CUNIA OOCOC	Clinian I Tain			Cultivated autologous human						cells transplanted back to the patient allow		
CLIN1-08686	Clinical Trial	1815	0 1811	limbal stem cells on human	5 8:	0 11 71	44.244.244	100		restoration and maintenance of a normal corneal		
Deng, UCLA	Stage Projects	IND	Corneal Blindness	amniotic membrane	Eye Disease	Cell Therapy	\$4,244,211	LSC	Autologous	surface.	Obtain an active IND	
BLOOD & CANCER	R THERAPEUTICS	5										
Blood Disorders					ı		ı			In	1	III
										Untreated alpha thalassemia major is almost		
										universally fatal in utero. Current treatment		
										requires in utero blood transfusions and monthly		
										blood transfusions for life or a bone marrow		
										transplant if a suitable donor is identified. The		
										proposed treatment is a maternal bone marrow		
CLIN2-09183	Clinical Trial		Almha Thalassamia	Maternal bone marrow derived						transplant in utero that takes advantage of	Cafatu and facaibility	
		Dh1			Dland Disaudou	Call Thorony	ć12 121 017	usc	Allegeneie	maternal-fetal immune tolerance, and may provide		
Mackenzie, UCSF	Stage Projects	Ph1	Major	HSC transplant in utero	Blood Disorder	Cell Therapy	\$12,131,817	HSC	Allogeneic	a definitive cure.	efficacy.	
										CGD prevents white blood cells from killing foreign		
										invaders. Patients have persistent, untreatable		
				Lentiviral vector modified						tissue infections. Affects 1:200,000 in US. Usually diagnosed before age 5, without treatment		
				autologous CD34+						, , , , , , , , , , , , , , , , , , ,	Primary: Safety and	
			X-linked Chronic	hematopoietic stem/progenitor		Genetically				children die before age 10. Project plan is transplantation of severe X-CGD patients that lack	Efficacy. Secondary:	
CLIN2-08231	Clinical Trial		Granulomatous	cells via transplantation &		Modified Cell				matched donors using gene-corrected autologous	Restoration of immune	
Kohn UCLA	Stage Projects	Ph 1/2	Disease.	engraftment	Blood Disorder	Therapy	\$7,402,549	HSC	Autologous	HSCT.	function	
KOIIII OCLA	Stage Frojects	1111/2	Disease.	engratthent	blood bisorder	тпетару	\$7,402,343	1150	Autologous	An inherited mutation in the hemoglobin gene	Tunction	
										causes red blood cells to "sickle" under conditions		
										of low oxygen. Affects 1:500 African-Americans and		
										is common in Hispanic-Americans. Median survival		
										is 42 years for males and 48 years for females.		
	Duane Roth			Autologous HSC, genetically						More than 80% of patients lack an HLA-identical		
	Disease Team			corrected ex vivo by lentiviral						sibling donor. Project plan is genetic correction of	Primary: Safety, feasibility.	
	Therapy			vector mediated addition of a		Genetically				adult bone marrow hematopoietic cells by adding a		
DR3-06945	Development			hemoglobin gene that blocks		Modified Cell				novel therapeutic hemoglobin gene that blocks	Recovery; RBC function;	
Kohn, UCLA	III	Ph 1	Sickle Cell Disease	sickling	Blood Disorder	Therapy	\$13,935,441	HSC	Autologous	sickling of the red blood cells.	Quality of life assessment	
,							, ,,,,,,,			In ADA-SCID allogeneic HSCTs from non-matched	,	
										sibling donors are a high risk procedure. Efficacy of	Primary: Safety.	
				Autologous HSC, genetically						chronic enzyme replacement therapy is uncertain	Secondary: Efficacy, gene	
			ADA-SCID (severe	corrected ex vivo by lentiviral		Genetically				in the long-term. Preliminary data indicates that	marking, immune	
CLIN2-09339	Clinical Trial		combined immune	vector mediated addition of		Modified Cell				OTL-101 may significantly improve outcomes	reconstitution.	
Kohn, UCLA	Stage Projects	Ph2	deficiency)	human ADA gene	Blood Disorder	Therapy	\$20,000,000	HSC	Autologous	compared to available therapies.	Registrational trial.	
İ										Catastrophic immunodeficiency disorder caused by	Primary: Safety and	
CLIN2-09504			X-SCID (X-linked	Autologous HSC, genetically		Genetically				mutation in IL2RG; Without a curative transplant-	feasibility. Secondary:	
Sorrentino, St.	Clinical Trial		severe combined	corrected ex vivo by lentiviral		Modified Cell				based therapy, X-SCID is lethal typically in first year		
Jude's	Stage Projects	Ph 1/2	immunodeficiency)	vector mediated correction	Blood Disorder	Therapy	\$11,924,780	HSC	Autologous	of life.	immune reconstitution	
			Conditioning									
			regimen for							Monoclonal antibody that targets CD117 and		
			allogeneic HSC							promotes engraftment of hematopoietic stem		
			transplantation for							cells. Could replace toxic conditioning regimens		
	Disease Team		SCID (Severe							and enable chemotherapy-free transplants.	Safety. Dosing. Efficacy -	
DR2A-05365	Therapy			MAb that depletes endogenous						Enabled donor cell HSC engraftment and cure of	HSC engraftment, immune	
Shizuru, Stanford	Development	IND, Ph 1	Immunodeficiency)	HSC	Blood Disorder	Biologic	\$19,068,382			disease in an animal model of SCID.	reconstitution.	
										Primary immune deficiency due to Artemis gene.		
										Most difficult to treat by allogeneic hematopoietic		
			ART-SCID (Artemis-							stem cell transplantation (HSCT) due to increased		
	Late Stage		deficient severe	Autologous HSC, genetically		Genetically				sensitivity to alkylating agents and radiation.		
CLIN1-08363,	Preclinical			corrected ex vivo by lentiviral		Modified Cell				Autologous gene modified HSCT transplantation to		
Puck, UCSF	Projects	IND	immunodeficiency)	vector mediated correction	Blood Disorder	Therapy	4,268,865	HSC	Autologous	overcome allogeneic stem cell transplant difficulty.	Obtain an active IND	
CLIN1-10084,	Late Stage			Autologous HSC, genetically		Genetically						
Porteus,	Preclinical			corrected ex vivo by CRISPR-		Modified Cell				Gene editing using CRISPR-Cas9 technology has the		
Stanford	Projects	IND	Sickle Cell Disease	mediated correction	Blood Disorder	Therapy	5,194,431	HSC	Autologous	potential to correct the sickle cell mutation.	Obtain an active IND	
HIV/AIDS												

Award							Funding	Therapeutic				Percent
Number, PI,					General Disease	General Class	(ICOC	Cell (for Cell				Time Into
Institution	Program	Trial Stage	Indication	Therapeutic	Area	of Approach	Approved)	Therapy)	Cell Source	Rationale	Project Goal	Award
				Autologous HSC transduced ex			,	. , ,				
				vivo with a lentiviral vector								
DR1-06893				engineered to express an		Genetically				Cal-1 increases the number of HIV-protected cells	Safety. Efficacy - slow	
Symonds,				shRNA against CCR5 & a fusion		Modified Cell				in the body. Uses shRNA to CCR5 and C46 to confer	disease progression,	
Calimmune	Disease Team I	Ph 1/2a	HIV/AIDS	inhibitor.	HIV/AIDS	Therapy	\$8,278,722	HSC	Autologous	cellular resistance to HIV infection.	mitigate need for ART.	
				Gene modified HSCs via a						Lentiviral vector encodes a triple combination of		
				lentiviral vector that encodes a						HIV-resistance genes and a pre-selective marker.		
				triple combination of HIV-		Genetically				Vector transduced CD34+ cells will safely engraft,	Safety. Efficacy - immune	
CTS1-08231	Clinical Trial			resistance genes and a tCD25		Modified Cell				divide and differentiate in vivo into mature myeloid	·	
Abedi, UC Davis	Stage Projects	Ph 1	HIV/AIDS	pre-selective marker	HIV/AIDS	Therapy	\$7,402,549	HSC	Autologous	and lymphoid cells.	and HIV status.	
										Autologous hematopoietic stem cells gene edited		
CD24 0752C						Compatible				ex vivo to eliminate expression of HIV entry co-		
SP3A-07536	Chunhania			Autologous USCs sousticelly		Genetically				receptor CCR5. Cells carrying disrupted CCR5	Cafat. Efficaci.	
Zaia, City of	Strategic	Dh 1		Autologous HSCs genetically	HIV/AIDC	Modified Cell	¢E E02 420	HSC	Autologous	provide a renewable, long-lasting source of HIV-1	Safety. Efficacy -	
Hope Hematologic Can	Partnership III	Ph 1	HIV/AIDS	modified to disrupt CCR5	HIV/AIDS	Therapy	\$5,583,438	HSC	Autologous	resistant immune cells.	engraftment.	
Hematologic Can	l			I		I	I	I	1	Cancer is a leading cause of death in CA. Many		
										cancers resist current therapies due to therapy-		
										resistant cancer stem cells (CSCs). Discovered a	Safety. Dosing. Follow on	
	Duane Roth									protein, ROR1, present on CSCs but not on normal	trials will include other	
	Disease Team									healthy cells. Developed an antibody,	cancers and will test	
	Therapy			Monoclonal antibody (anti-						cirmtuzumab, that is specific for ROR1. Project plan		
DR3-06924	Development			ROR1) targeting CLL cancer	Hematologic					is to treat chronic lymphocytic leukemia with	combination with other	
Kipps, UCSD	III	Ph 1	CLL	stem cells	Malignancy	Biologic	\$4,179,600			cirmtuzumab.	anti-cancer therapies.	
										Cancer is a leading cause of death in CA. Many		
										cancers resist current therapies due to therapy-		
										resistant cancer stem cells (CSCs). Discovered a		
										protein, ROR1, present on CSCs but not on normal		
										healthy cells. Developed an antibody,		
										cirmtuzumab, that is specific for ROR1. Project plan		
				Monoclonal antibody (anti-						is to treat chronic lymphocytic leukemia or mantle		
CLIN2-10192	Clinical Trial			ROR1), combined with tyrosine	Hematologic					cell carcinoma with cirmtuzumab in combination	Evaluate dosing and	
Kipps, UCSD	Stage Projects	Ph 1b/2a	B Cell Cancers	kinase inhibitor Ibrutinib	Malignancy	Biologic	\$18,292,674			with ibrutinib.	complete response rate.	
										CD34+ hematopoietic Stem and progenitor cells		
										engraft into the bone marrow of patients,		
										rebuilding a new blood and immune system after		
										appropriate preparation called myeloablation. The		
	1							Expanded CD34+		endothelial cells used in the co-culture are thought		
	1			Matched cord blood derived				stem and progenitor cells		to aid the engraftment of the stem and progenitor cells into the bone marrow via secretion of		
	1		Hematologic	hematopoietic stem and				from cord blood		angiocrine factors. The remainder of the cord		
CLIN1-08342	1		•	progenitor cells expanded by co-				and gene-		blood cells in the cell product also aid in the		
Davis, Angiocrine	Clinical Trial		including leukemia	culture with genetically	Hematologic			modified		engraftment as well as provide anti-viral and anti-		
Bioscience	Stage Projects	IND	and lymphoma	modified endothelial cells.	Malignancies	Cell Therapy	\$3,797,117	endothelial cells	Allogeneic	bacterial effects after transplantation.	Obtain an active IND	
2.000lenee	_ tage . rojects		3.1.4 1,piioilia	I I I I I I I I I I I I I I I I I	ag.iuncies	cen merupy	70,.01,111		,ogenere	CD34+ hematopoietic Stem and progenitor cells	ann an active nep	
	1									engraft into the bone marrow of patients,		
	1									rebuilding a new blood and immune system after		
	1									appropriate preparation called myeloablation. The		
	1							Expanded CD34+		endothelial cells used in the co-culture are thought		
	1							stem and		to aid the engraftment of the stem and progenitor		
	1			Matched cord blood derived				progenitor cells		cells into the bone marrow via secretion of		
CLIN2-10386	1		Hematologic	hematopoietic stem and				from cord blood		angiocrine factors. The remainder of the cord		
Finnegan,	1		malignancies	progenitor cells expanded by co-				and gene-		blood cells in the cell product also aid in the		
Angiocrine	Clinical Trial		_	culture with genetically	Hematologic			modified		engraftment as well as provide anti-viral and anti-		
Bioscience	Stage Projects	Ph 1b	and lymphoma	modified endothelial cells cells.	Malignancies	Cell Therapy	\$5,000,000	endothelial cells	Allogeneic	bacterial effects after transplantation.	Safety.	1

Award			1	I			Funding	Therapeutic			October Zotri, z	Percent
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Number, PI,	_				General Disease		(ICOC	Cell (for Cell				Time Into
Institution	Program	Trial Stage	Indication	Therapeutic	Area	of Approach	Approved)	Therapy)	Cell Source	Rationale	Project Goal	Award
										ADCs are intended to target and kill only the target	Obtain an active IND	
										cancer cells and spare healthy cells. ADCs are		
										composed of an antibody linked to a cytotoxic		
										payload or drug. After the ADC binds to the target		
										cell and is internalized, the cytotoxic drug is		
										released and kills the cancer cell. CLL1 is highly		
CLIN1-09776	Late Stage									expressed on leukemia stem cells but not on		
Junutula,	Preclinical			Anti-CLL1 antibody linked to a	Hematologic	Antibody-drug				normal cells. Binding of the anti-CLL1 ADC results in		
Cellerant	Projects	IND	AML	DNA binding payload.	Malignancy	conjugate (ADC)	\$6,863,755			targeted killing of leukemia stem cells.		
CLIN2-09574	Clinical Trial			O , · · · ,	Hematologic	, , , ,	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			Using cord blood transplant to overcome the		
		Ph 2	AML	Umbilical cord blood stem cells.		Cell Therapy	¢¢ 022 100	Cord blood	Allegeneie	neutropenia caused by chemotherapy for AML.		
Delaney, Nohla	Stage Projects	PILZ	AIVIL	Ombilical cord blood stem cells.	Malignancy	Сен тпетару	\$6,922,109	Cora biooa	Allogeneic			
										CD47 is overexpressed on cancer and cancer stem		
										cells and prevents their elimination by phagocytic		
										macrophages by delivering a potent "don't eat me"		
										signal. Hu5F9-G4 is a humanized monoclonal		
										antibody (mAb) that binds to CD47 and blocks its		
CLIN2-10144	Clinical Trial			Anti-CD47 monoclonal antibody	Hematologic					interaction with its receptor, thereby enabling		
		DI 41			_	B: 1 ·	ÅT 000 000					
Chao, 47Inc	Stage Projects	Ph 1b	AML	with azacitidine	Malignancy	Biologic	\$5,000,000			phagocytosis of cancer cells.	l	
Solid Cancers	ı			1	ı	ı	1	ı			ı	
]		ĺ]			There are few options for patients whose cancers	Primary: Safety and	
]		Advanced tumors]			have metastasized due to resistance to current	feasibility. Secondary:	
	Disease Team		(Synovial Sarcoma,	Autologous HSCs and T cells		Genetically]			therapies. Engineering of patient's own blood-	Persistence of gene-	
DR2A-05309	Therapy		Melanoma,	genetically modified to express		Modified Cell				forming stem cells to produce a continual supply of	_	
Ribas, UCLA	Development	IND, Ph 1	Ovarian)	an anti-tumor T cell receptor.	Solid Tumor	Therapy	\$19,999,563	HSC	Autologous	the immune system cell to attack cancer.	immune cells	
NIDAS, UCLA	Development	IND, FILL	Ovarially	an anti-tumor i cen receptor.	John Turrior	Петару	\$15,555,505	пзс	Autologous		illillidile cells	
										Solid tumors are the most prevalent form of		
										cancer, and are a major cause of death worldwide.		
	Duane Roth									The small molecule being developed inhibits the	Determination of	
	Disease Team			Small molecule mitotic inhibitor						activity of a protein required in tumor cell lines and	maximum tolerated dose	
	Therapy			targeting serine/threonine						cancer stem cells (CSC). It is hypothesized that	and recommended Phase	
DR3-07067	Development			kinase to eliminate both tumor						inhibiting the CSC can prevent tumor regrowth	2 dose. Safety. PK. Efficacy	
	-	DI- 1	Solid Tumor	cells and cancer stem cells	Callel Townson	Constitution of the contra	¢c 024 247			after treatment.		
Slamon, UCLA	III	Ph 1	Solia Tumor	cells and cancer stem cells	Solid Tumor	Small Molecule	\$6,924,317				in solid cancers.	
										CD47 is overexpressed on cancer and cancer stem		
										cells and prevents their elimination by phagocytic		
										macrophages by delivering a potent "don't eat me"		
										signal. Hu5F9-G4 is a humanized monoclonal		
										antibody (mAb) that binds to CD47 and blocks its		
										interaction with its receptor, thereby enabling		
										phagocytosis of cancer cells. Anti-CD47 is highly		
											Cofety Desire Efficient	
										synergistic in combination with other anti-cancer	Safety. Dosing. Efficacy -	
CLIN2-09577	Clinical Trial			Anti-CD47 monoclonal antibody						therapies including tumor-targeting mAbs such as	objective response rate	
Chao, 47Inc	Stage Projects	Ph1b/2	Solid Tumor	+ cetuximab	Solid Tumor	Biologic	\$10,234,048	Ab	_	cetuximab.	(ORR)	
						Genetically						
CLIN2-10248	Clinical Trial		ĺ	T cells engineered to target		Modified Cell]					
Brown, COH	Stage Projects	Ph 1	Malignant Glioma	-	Solid Tumor	Therapy	\$12,753,854	CAR-T	Autologous			
			Wangilanc Giloma	carreer steril cens	Sona ramor	тистиру	ψ12,733,03·	G/ 11 ()	/ tatologous			
ORGAN SYSTEMS	THERAPEUTICS)										
Bone Disorders			•				•		_		1	
]		ĺ]			Femoral head osteonecrosis (aka avascular		
]		ĺ]			necrosis) is a disease caused by loss of blood supply		
]		ĺ]			to the bone, leading to bone cell death, end stage		
]		ĺ				Ì			hip arthritis and total hip replacement. There is an		
			1									
]		ĺ				Ì			unmet need for treatment of this disease, that	L	
			1							affects individuals at prime of life (peak age 35	Safety, tolerability.	
	Ì		İ	1			l		1	years). This small molecule therapeutic recruits	Determine PK. Determine	
]		ĺ	Synthetic molecule, LLP2A-Ale,			Ì			bone forming cells to site of damage, where they	PD effects on bone	
	Disease Team		ĺ	to enhance homing of			Ì			serve the dual function of laying down new bone,	turnover, biomarkers.	
DB34 0E303			İ	_			l		1			
DR2A-05302	Therapy	DI 6 "	l	endogenous bone marrow			440.000.00			and stimulating revascularization to prevent	Determine	
Lane, UC Davis	Development	Ph 1 a/b	Osteonecrosis	MSCs to bone surface	Bone Disorder	Small Molecule	\$19,999,867			further bone cell death.	immunogenicity.	
Cartilage Disorde	ers											
CLIN1-09472												
Wang, Cellular	Late Stage		ĺ				Ì					
Biomedicine	Preclinical		ĺ	Allogeneic adipose-derived			Ì					
Group	Projects	IND	Osteoarthritis	MSCs	Cartilage Disorder	Cell Therapy	\$2,291,976	MSC	Allogeneic		Obtain an active IND	
огоир	riojects	IND	Osteodi tili itiS	111303	carmage Disorder	cen merapy	72,231,310	IVIJC	Allogerieit		Optain an active IND	

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Award					C15'	C	Funding	Therapeutic				Percent
Number, PI,	D	T.::-1.0:	to die er	There are	General Disease		(ICOC	Cell (for Cell	C-11 C	B-sti.	Dunia de C	Time Into
Institution	Program	Trial Stage	Indication	Therapeutic	Area	of Approach	Approved)	Therapy)	Cell Source	Rationale	Project Goal	Award
				Small molecule injected intra-								
				articularly that promotes								
CLIN1-08309			Osteoarthritis,	resident cartilage mesenchymal stem cell differentiation into								
Schultz, Calibr	CLIN1	IND	cartilage injuries	chondrocytes	Cartilage Disorder	Small Molecule	\$1 667 832				File an IND	
Cardiovascular &			car thage injuries	chondrocytes	curtilage Disorder	Sman Wolcculc	\$1,007,03Z				THE UT IND	
											Primary: Determine	
											whether treatment is safe	
											and causes reduction in	
											cardiac scar size in patients	5
											with heart failure after a	
2224 25725			Heart dysfunction							Heart failure is a progressive disease with a high	heart attack. Secondary:	
DR2A-05735	Disease Team		after myocardial	Allaganaia saudiaan bana daninad	Condinuesculos					risk of mortality. Cardiosphere-derived cells (CDCs)	Assess for other structural or functional cardiac	
Smith, Capricor Inc.	Therapy Development	Ph 2	heart failure	Allogeneic cardiosphere derived cells	Cardiovascular Disease	Cell Therapy	\$19,782,136	CDC	Allogeneic	reduce scar size after heart attack in preclinical animal models and in a prior clinical trial.	benefits.	
IIIC.	Development	FILE	neart failure	cens	Disease	cell filerapy	\$13,762,130	СВС	Allogeneic	Pulmonary arterial hypertension (PAH) is a	bellelits.	
										progressive condition with no cure, survival is poor.		
										Cardiosphere-derived cells (CDCs) decrease wall		
										thickening of lung small blood vessels in preclinical	Primary: Safety.	
CLIN2-09444										studies. Improvement in lung blood vessels is	Secondary: Exploratory	
Lewis, Cedars-	Clinical Trial			Allogeneic cardiosphere derived						expected to reduce cardiac right ventricular	efficacy measures of right	
Sinai	Stage Projects	Ph1a/b	Hypertension	cells	Vascular Disease	Cell Therapy	\$7,354,772	CDC	Allogeneic	dysfunction.	ventricular function.	
										5.7 million Americans suffer from heart failure, and		
	Disease Team									the end stage 2 year survival rate is 50%. hESC-CM promote new blood vessel formation and improve	Obtain an active IND for a	
DR2A-05394	Therapy		Ischemic heart	Allogeneic hESC-derived	Cardiovascular					cardiac function in preclinical models of heart	first-in-human trial in heart	
Wu, Stanford	Development	IND	failure	cardiomyocytes	Disease	Cell Therapy	\$19,060,330	CM	Allogeneic	failure.	failure patients.	
Diabetes & Comp		bolic				. ,						
										Diabetes mellitus affects 370 million people		
										worldwide. Disproportionately affects certain		
										minority groups and the elderly. Current therapy is		
										self-administration of insulin. Diabetes costs in CA		
										are tens of billions of dollars each year. Directed differentiation of embryonic stem cells to		
										pancreatic precursor cells. Project plan is		
				Allogeneic hESC-derived						transplantation of pancreatic precursor cells that		
AP1-08039				pancreatic cell progenitors in a				Pancreatic		generate functional islet tissue in vivo that can		
Foyt, ViaCyte	Accelerated	Comparabili		device implanted		Cell Therapy,		endocrine		respond to insulin levels in a more physiological	Primary: Safety.	
Inc.	Pathway I	ty Trial	Diabetes: Type 1	subcutaneously	Endocrine Disorder	Combination	\$16,603,160	progenitor	Allogeneic	manner than direct insulin replacement.	Secondary: Efficacy.	
										Children with T1D face lifelong struggles with		
										glycemic control and, despite careful management,		
										an increased risk of severe complications. No		
										therapy that maintains or restores pancreatic beta islet cell function is currently		
										approved. Evidence indicates that		
CLIN2-09730										regulatory T-cells (T-regs) maintain immune		
Losordo,	Clinical Trial			Autologous ex vivo expanded						balance at least in part by control of differentiation	Primary: Safety.	
Caladrius	Stage Projects	Ph 2	Diabetes: Type 1	polyclonal regulatory T cells	Endocrine Disorder	Cell Therapy	\$12,211,255	T-reg	Autologous	of multipotent progenitor/stem cells.	Secondary: Efficacy.	
								-		There are over 100,000 people in the US with type		
										1 diabetes so severe that they are at constant risk		
				hESC-derived pancreatic						of hospitalization and/or death. Within months		
CUNA COSTA				progenitor cells delivered in a				D ::		after administration, this product could provide a		
CLIN1-08671,	Clinia - LT-1-1			device that allows direct		Call Theren		Pancreatic		source of insulin producing beta cells to restore	Ohtoin on outive IND -	
D'Amour,	Clinical Trial Stage Projects	IND	Diabetes: Type 1	vascularization of the cell	Endocrine Dicarda-	Cell Therapy, Combination	\$3,984,164	endocrine	Allogonois	those patients' blood sugar to normal healthy levels and save their lives.	Obtain an active IND and trial start up	
Viacyte	stage Projects	טאוו	Diabetes: Type 1	therapy	Endocrine Disorder	Combination	23,384,104	progenitor	Allogeneic	There are over 100,000 people in the US with type	triai start up	
										1 diabetes so severe that they are at constant risk		
	1			hESC-derived pancreatic						of hospitalization and/or death. Within months		
				progenitor cells delivered in a						after administration, this product could provide a		
				device that allows direct				Pancreatic		source of insulin producing beta cells to restore		
CLIN2-09672,	Clinical Trial			vascularization of the cell		Cell Therapy,		endocrine		those patients' blood sugar to normal healthy		
Foyt, Viacyte	Stage Projects	Ph 1/2	Diabetes: Type 1	therapy	Endocrine Disorder	Combination	\$20,000,000	progenitor	Allogeneic	levels and save their lives.		

Award							Funding	Therapeutic				Percent
Number, PI,					General Disease		(ICOC	Cell (for Cell				Time Into
Institution	Program	Trial Stage	Indication	Therapeutic	Area	of Approach	Approved)	Therapy)	Cell Source	Rationale	Project Goal	Award
										Cystinosis is caused by a genetic mutation that		
										allows an amino acid, cystine, to build up in and		
										damage the kidneys, eyes, liver, muscles, pancreas		
										and brain of children and adults. Current therapy		
										only delays progression of the disease, has severe		
										side effects and people taking it still require kidney		
										transplants, and develop diabetes, neuromuscular		
										disorders and hypothyroidism. The goal is to take		
				Ex vivo transduced autologous						blood stem cells from people with cystinosis,		
				human CD34+ hematopoietic		Genetically				genetically-modify them to remove the mutation,		
Clin1-09230	Clinical Trial			stem cells for treatment of		Modified Cell				then return them to the patient to create a new,		
Cherqui, UCSD	Stage Projects	IND	Cystinosis	cystinosis	Cystinosis	Therapy	\$ 5,273,189	HSC	Autologous	healthy, blood system free of the disease.	Obtain an active IND	
Skeletal Muscle I	Disorders				I				T T	I	I	
										Heart failure is a leading cause of death for	Primary: Safety and	
										Duchenne muscular dystrophy patients.	tolerability in DMD	
			Duchenne							Cardiosphere-derived cells (CDCs) decrease	patients. Secondary:	
CLIN2-08334			muscular							myocardial fibrosis, improve cardiac function and	Structural or functional	
Ascheim,	Clinical Trial		dystrophy	Allogeneic cardiosphere derived	Skeletal Muscle					induce regeneration of heart muscle in preclinical	cardiac benefits, quality of	
Capricor, Inc.	Stage Projects	Ph 2	cardiomyopathy	cells	Disorder	Cell Therapy	\$3,376,259	CDC	Allogeneic	models of DMD.	life improvements.	
Other Disorders		1		I	I				T	la		
				A Human Acellular Vessel in						Synthetic vascular access grafts for hemodialysis in		
				Patients Needing Renal						kidney patients are associated with thrombosis,	Primary: Safety and	
0.11.0				Replacement Therapy: A						infection and abandonment. Human Acellular	tolerability, rate of	
CLIN2-08938,	Clinia di Taial			Comparison with ePTFE Grafts						Vessel (HAV) is made of extracellular matrix from	patency of the graft and	
Lawson,	Clinical Trial	DI- 2	Daniel dieberte	as Conduits for Hemodialysis	For the color of Disconding	D	¢0.000.530		A.II	human smooth muscle cells, similar in composition		
Humacyte, Inc.	Stage Projects	Ph 3	Renal dialysis	(HUMANITY)	Endocrine Disorder	Device	\$9,999,528			and structure to native tissue.	needed to restore patency.	
										Synthetic vascular access grafts for hemodialysis in		
										kidney patients are associated with thrombosis,		
CLINIA COCCO				A Human Acellular Vessel in						infection and abandonment. Human Acellular Vessel (HAV) is made of extracellular matrix from		
CLIN2-09688,	Clinical Trial									· · · ·	A Campanianaith A\/	
Lawson,		Dh 2	Bonal dialusis	Patients Needing Renal	Endocrino Dicardor	Dovice	¢14 002 005		Allogons:	human smooth muscle cells, similar in composition	Fistula	
Humacyte, Inc.	Stage Projects	Ph 3	Renal dialysis	Replacement Therapy.	Endocrine Disorder	Device	\$14,082,865		Allogeneic	and structure to native tissue.	ristuid	
										Unmet medical need for allogeneic kidney		
										transplants. Need to eliminate chronic		
										rejection/allograft nephropathy that causes		
CLINI2 00420				Damas CD241 and CD21 Tability						gradual loss of kidney (50% of graft loss by 12-15	Duine and Cofety	
CLIN2-09439	Clinia di Tairi		T	Donor CD34+ and CD3+ T cells						years in HLA mismatched recipients). Eliminate the		
Strober,	Clinical Trial		Transplant	for immune tolerance to HLA	Immune tolerance,	0.11.71	45.000.00			lifelong need for anti-rejection drugs that have	Secondary: Preliminary	
Stanford	Stage Projects	Ph 1	tolerance	mismatched kidney donors.	transplant	Cell Therapy	\$5,069,674	HSC	Allogeneic	numerous cumulative side effects.	efficacy.	